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CALGENE INC

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C12n-15

cDNA sequence coding for plant acyl carrier protein - used for detection and prepn. of expression constructs esp. for prodn. of seed

R(AT BE CH DE ES FR GB GR IT LI LU NL SE)

A cDNA sequence coding for plant acyl carrier protein (ACP)

The gene can be obtd. from plants such as spinach, Brassica campestris or B.napus.

Sequences coding for plant ACP can be used as probes for detecting the presence of ACP genes, for screening libraries from plants and bacteria and for use in assays for detecting the presence of ACP genes. The coding sequences can be used in the prepn. of expression constructs, esp. with a transcription initiation region which is functional in a plant nil regulated so as to provide expression in seed. In this armner, the prodn. of seed oil can be enhanced and, as appropriate, the fatty acid compsn. modulated.

C(4-B4A1, 4-B4A4, 11-A, 12-K4D) D(5-H12)

PRODUCTION

Among various ways in which the gene may be obtd., a library may be prepd., either genomic or cDNA. Probes may be prepd. based on the amino acid sequence of the ACP.

Both prokaryotic and eukaryotic polyclonal antibodies may be used for isolating ACPs from a particular source and may be further used to isolate ACPs from other plant sources. The ACP may then be sequenced and probes designed based on the peptide sequence.

EXAMPLE

Total RNA was extd. from young spinach leaves in 4M guanidine thiocyanate buffer. Total RNA was subjected to oligo(dT)-cellulose column chromatography 2 times to yield poly(A)+RNA. A cDNA library was constructed in pUC13-Cm. The annealed cDNA was transformed into competent E.coli JM83 cells and spread onto LB agar plates contg. 50µg/ml chloramphenicol and 0.005% X-Gal.

A total of 8000 cDNA clones were screened by performing Southern blots. A 5'-end-labelled synthetic oligonucleotide (ACPP4) that is at least 66% homologous with a 16 amino

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Best Available Cor

acid region of spinach ACP-I was used as an ACP probe. The positive clone, pCGNISOL, was subcloned into pUC118 and pUC119. Single-stranded DNA template was prepd. and DNA sequence was detd. using the Sanger dideoxy technique.(27pp 1703LDDwgNo0/0). (E) ISR:- No Search Report.	
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Acyl carrier protein- DNA sequence and synthesis.

DNA sequences are provided coding for acyl carrier protein, which sequence can be used for production of acyl carrier protein as an end product or in plant seed to enhance seed oil production. A regulated promoter is provided which substantially limits expression of the acyl carrier protein to seed tissue.

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Description

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ACYL CARRIER PROTEIN - DNA SEQUENCE AND SYNTHESIS

FIELD OF THE INVENTION

Acyl carrier protein is expressed under conditions where the protein may be isolated for in vitro use or the protein is intracellularly translocated to a chloroplast or related organelle for modification of fatty acid production in vivo. Constructs are provided which allow for expression of acyl carrier protein in seed tissue using a seed specific promoter.

BACKGROUND OF THE INVENTION

Plants provide a rich source of a variety of products which find use in foods, as raw materials, and as finished products. Vegetable fatty acids find extensive use for a wide variety of commercial purposes, being used as vegetable oils for cooking, as lubricants, in alkyd resins, as specialty chemicals, and the like. For the most part, the plant fatty acids tend to be of 18 carbon atoms, there usually being only a minor level of fatty acids having fewer than 16 carbon atoms. For many purposes, it would be desirable to have fatty acids in the range of 8 to 14 carbon atoms. There is, therefore, substantial interest in developing methods for producing vegetable oils where there is a substantial proportion of the total fatty acids of 14 carbon atoms or fewer.

To achieve this purpose, it will be necessary to modify the constituent members of the metabolic pathway resulting in the formation of fatty acids and their elongation to higher fatty acids. Toward this purpose, it will be necessary to be able to produce one or more components along the fatty acid metabolic chain which modify the course of the plant metabolism. In addition, there may be significant commercial applications for individual components of the fatty acid metabolic pathway.

BRIEF DESCRIPTION OF THE RELEVANT LITERATURE

Kuo and Ohlrogge, Archives of Biochem. and Biophys. (1984) 234:290-296, describe the primary structure of spinach acyl carrier protein. Ohlrogge and Kuo, J. Biol. Chem. (1985) 260:8032-8037 report the existence of different isoforms of acyl carrier protein expressed differently in different tissues. Crouch et al., J. Mol. Appl. Genet. (1983) 2:273-283, report the synthesis of cDNA coding for napin protein.

SUMMARY OF THE INVENTION

DNA constructs are provided which provide for expression of plant acyl carrier protein. Particular constructs are produced which employ transcription initiation regions resulting in expression in plant embryos during seed maturation. The composition and amount of fatty acid can be modulated by modifying the constituents in the chloroplast or related organelle involved in a metabolic pathway in the production of fatty acids.

DESCRIPTION OF SPECIFIC EMBODIMENTS

Methods and compositions are provided for production of acyl carrier protein as an end product for use in vitro or in conjunction with seed formation of plants to provide for modified expression of fatty acids in vivo. Towards this end, DNA constructs are prepared, where the sequence encoding plant acyl carrier protein is joined to transcriptional initiation and termination regulatory regions, which are func tional in a predetermined host for expression of the acyl carrier protein.

The expression constructs provide in the 5'-3' direction of transcription, a transcriptional initiation regulatory region, either constitutive or regulated, an open reading frame coding for at least a functional portion of the acyl carrier protein, desirably including a transit peptide sequence providing for translocation to the chloroplast for in vivo use, and a transcriptional termination regulatory region functional in the appropriate host.

Depending upon the host, the regulatory regions will vary. For expression in a prokaryotic or eukaryotic microorganism, particularly unicellular, host, a wide variety of constitutive or regulatable, promoters may be employed. In these instances, the primary purpose for the preparation of the acyl carrier protein is the use of the acyl carrier protein for in vitro applications.

For the most part, the constructs will involve regulatory regions functional in plants which provide for enhanced production of acyl carrier protein for enhanced and/or modification of the fatty acid composition.

The coding sequence which is employed may be derived from natural sources, synthesized, or combinations thereof. To obtain the gene from a natural source, any of a variety of plants or bacteria may be used as the source of the gene. Plants include spinach, Brassica, e.g. campestris or napus. coconut, cotton, safflower, sunflower, Cuphea etc. Among various ways in which the gene may be obtained, a library may be prepared, either genomic or cDNA. Probes may be prepared based on the amino acid sequence of the acyl carrier protein. Since it is found that there is a substantial immunological cross-reactivity between acyl carrier proteins from different sources, both prokaryotic and eukaryotic polyclonal antibodies may be employed for isolating acyl carrier proteins from a particular source and may be further used to isolate acyl carrier proteins from other plant sources. The acyl carrier protein may then be sequenced in whole or in part, and probes designed based on the peptide sequence. Where only a partial DNA sequence is obtained, the partial

sequence m been obtain Once the sequence in example, a r in vitro muta Thus, trans sequence. In modified to a provide for a may be furt restriction s The acyl c ACP-I and A in other pla Of particu sequence.

sequence may be satisfactory or the gene may be walked, so as to ensure that the entire coding sequence has been obtained.

Once the desired sequence has been obtained, it may be manipulated in a variety of ways. Where the sequence involves non-coding flanking regions, the flanking regions may be subjected to resection, using, for example, a nuclease such as Bal 31, restriction with a restriction endonuclease, or modification by employing in vitro mutagenesis, primer repair, or other methods for introducing mutations or lesions into the sequence. Thus, transitions, tranversions, deletions, and insertions may be performed on the naturally occurring sequence. In addition, all or a portion of the sequence may be synthesized, where one or more codons may be modified to provide for a modified amino acid sequence, or one or more codon mutations may be introduced to provide for a convenient restriction site or other purpose involved with construction or expression. The gene may be further modified by employing synthetic adaptors, linkers to introduce one or more convenient restriction sites, or the like.

The acyl carrier protein may be any one of the isozymes which may be found in a particular host, such as ACP-I and ACP-II as designated in Ohlrogge and Kuo. <u>supra</u>, as found in spinach, or their analogues as found in other plant hosts.

Of particular interest is the spinach acyl carrier protein, more particularly ACP-I which has the following sequence.

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Spinach ACP-I

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The sequences of interest will generally have at least 300 bp usually at least about 360 bp, and desirably 411 base pairs, where the entire coding sequence includes 411 base pairs. In addition, there may be both 5' and 3'

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non-coding flanking regions which may extend from 1 bp to 200 bp or more from the 5' or 3' terminus of the coding region, there usually being fewer than about 100 bp, preferably fewer than about 10 bp 5' of the initiation codon, of the naturally occurring non-coding flanking region.

The open reading frame, coding for the acyl carrier protein or functional fragment thereof will be joined at its 5' end to a transcriptional initiation regulatory region. Numerous transcriptional initiation regulatory regions are available, which provide for a wide variety of constitutive or regulatable, e.g. inducible, transcription of the structural gene. Depending upon the host, transcriptional initiation regulatory regions may include regions from structural genes from viral, plasmid or, chromosomal genes, or the like. Among transcriptional initiation regions which have been described are regions from bacterial and yeast hosts, such as \underline{E} . coli, Bacillus subtilis, Saccharomyces cerevisiae, including genes such as β -galactosidase, lambda left and right promoters, glycolytic enzyme promoters, and the like. Among transcriptional initiation regions used for plants are such regions associated with the structural genes for nopaline, octopine, mannopine, ribulose-1,3-bisphosphate carboxylase, the large and small subunits, the full length promoter from cauliflower mosaic virus, napin, phaseolin, etc.

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Of particular interest are those promoters which are regulated during seed maturation, particularly those synthesized in cotyledons of the embryo. These regulatory regions include regulatory regions of genes such as napin, phaseolin and glycinin. Napin regulatory regions of particular interest are from <u>Brassica</u> species, more particulary <u>campestris</u> and <u>napus</u>. The regulatory region will generally be at least about 150 bp and not more than about 3500 bp usually not more than about 2500 bp, and desirably not more than about 1000 bp. The napin gene has been described, Crouch, et al., <u>supra</u>, although there has been no disclosure of the regulatory region, nor the use of the regulatory region for an heterologous gene.

The transcriptional initiation regulatory region and coding region may be joined directly, where there are convenient restriction sites for the two regions or such restriction site(s) have been introduced, or, as appropriate, by means of synthetic adaptors or linkers. A number of regulatory regions are available as plasmids where the initiation and termination regulatory regions are separated by a polylinker, so that a number of restriction sites are available for insertion of the structural gene. These expression constructs are mostly available for microorganism hosts.

While a number of transcriptional initiation and termination regions functional in plants have been isolated, particularly from genes on the Ti- and Ri-plasmids, these regions have not achieved the level of readily available constructs including polylinkers, markers, replication systems and the like. Furthermore, for the present invention there is primary interest for expression to be regulated so that transcription is initiated in seed. For this purpose, a gene such as the napin gene is of substantial interest.

A napin regulatory region can be obtained by employing a probe which comprises a sequence adjacent the 3'- or 5'-terminus or intermediate coding sequence of the structural gene for screening a genomic library of the napin host, in the case of Crouch, 1983, <u>supra</u>, the rapeseed host. By identifying fragments which hybridize under stringent conditions with the probe, <u>fragments</u> having the napin structural gene may be identified. Potential regulatory sequences 5' of the napin structural gene can be identified by restriction mapping and DNA sequence analysis. These sequences can be manipulated under various conditions to remove in whole or in part the codons coding for napin, leaving the uncoded 5' region free or substantially free of the napin coding region. In some instances, it may be desirable to remove a short non-coding region adjacent the initiation codon, usually fewer than about 20 bp, more usually fewer than about 10 bp. For further details, the experimental section should be consulted.

After joining of the open reading frame for the acyl carrier protein structural gene and the transcriptional initiation regulatory region, a functional transcriptional termination regulatory region may be present, which has been included as a result of the method for construction or one may be introduced. The termination region may be from the same structural gene as the initiation region, acyl carrier protein gene, as is convenient. The termination region will usually include a terminator and sequence coding for polyadenylation.

The gene may naturally include or be modified by introducing a signal sequence for intracellular translocation, particularly to the leucoplast in seed or the chloroplast in other plant cells.

In developing the expression construct, the various components of the expression construct or fragments thereof will normally be inserted into a convenient cloning vector which is capable of replication in a bacterial host, e.g., <u>E. coli.</u> Numerous vectors exist that have been described in the literature. After each cloning, the plasmid may be isolated and subjected to further manipulation, such as restriction, insertion of new fragments, ligation, deletion, resection, insertion, in vitro mutagenesis or primer repair, so as to tailor the components to the desired sequence. Once the construct has been completed, it may then be transferred to an appropriate vector for further manipulation in accordance with the manner of transformation of the plant cell.

Normally, included with the expression construct will be a structural gene having the necessary regulatory regions for expression in a host and providing for selection of transformant cells. The gene may provide for resistance to a cytotoxic agent, e.g. antibiotic, heavy metal, toxin, etc., complementation providing phototrophy to an auxotrophic host, viral immunity or the like. Depending upon the number of different host species the expression construct or components thereof are introduced, one or more markers may be employed, where different conditions for selection are used for the different hosts.

The manner in which the construct is introduced into the plant host is not critical to this invention. Any method which provides for efficient transformation may be employed. Various methods include the use of Tior Ri-plasmids, microinjection, electroporation, liposome fusion, or the like. In many instances, it will be

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desirable to have the construct bordered on one or both sides by T-DNA, particularly having the left and right borders, more particularly the right border. This is particularly useful when the construct uses <u>A. tumefaciens</u> or <u>A. rhizogenes</u> as a mode for transformation, although the T-DNA borders may find use with other modes of transformation.

Where the Agrobacterium is used for plant cell transformation, a vector may be used which may be introduced into the Agrobacterium host for homologous recombination with T-DNA of the Ti- or Ri-plasmid present in the Agrobacterium host. The Ti- or Ri- plasmid containing the T-DNA for recombination may be armed (capable of causing gall formation) or disarmed (incapable of causing gall formation), the latter being permissible, so long as the vir genes are present in the transformed host. The armed plasmid can give a mixture of normal plant cell and gall.

In some instances where Agrobacterium is used as the vehicle for transforming plant cells, the expression construct bordered by the T-DNA border(s) will be inserted into a broad host spectrum vector, there being broad host spectrum vectors described in the literature. Commonly used in pRK2 or derivatives thereof. See, for example, Ditta et al. (1980) PNAS USA, 77:7347-7351 and EPA 0 120 515, which are incorporated herein by reference. Included with the expression construct and the T-DNA will be one or more markers, which allow for selection of transformed Agrobacterium and transformed plant cells. A number of markers have been developed for use with plant cells, such as resistance to chloramphenicol, the aminoglycoside G418, hygromycin, or the like. The particular marker employed is not essential to this invention, one or another marker being preferred depending on the particular host and the manner of construction.

The expression constructs may be employed with a wide variety of plant life, particularly plant life involved in the production of vegetable oils. These plants include <u>Brassica</u> e.g. <u>napus</u> and <u>campestris</u>, sunflower, safflower, cotton, Cuphea, soybean, and corn.

For transformation of plant cells using Agrobacterium, explants may be combined and incubated with the transformed Agrobacterium for sufficient time for transformation, the bacteria killed, and the plant cells cultured in an appropriate selective medium. Once callus forms, shoot formation can be encouraged by employing the appropriate plant hormones in accordance with known methods and the shoots transferred to rooting medium for regeneration of plants. The plants may then be grown to seed and the seed used to establish repetitive generations and for isolation of vegetable oils.

The DNA sequences can also be used as probes for searching for acyl carrier proteins in hosts other than the host from which the gene was derived. In addition, the acyl carrier protein produced in accordance with the subject invention can be used in preparing antibodies for assays for detecting acyl carrier protein. The acyl carrier protein can also be used in conjunction with chloroplast lysates to enhance the production and/or modify the composition of the fatty acids prepared in vitro. The acyl carrier protein can also be used for studying the mechanism of fatty acid formation in plants and bacteria.

The following examples are offered by way of illustration and not by way of limitation.

EXPERIMENTAL

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MATERIALS AND METHODS

Cloning Vectors

Cloning vectors used include the pUC vectors, pUC8 and pUC9 (Vieira and Messing, 1982) Gene. 19:259-268; pUC18 and pUC19 (Norrander et al., 1983) Gene, 26:101-106, Yanisch-Perron et al., (1985) Gene. 33:103-119, analogous vectors exchanging chloramphenicol resistance (CAM) as a marker for the ampicillin resistance of the pUC plasmids described above (pUC-CAM [pUC12-Cm, pUC13-Cm] Buckley, K., Ph.D. Thesis, U.C.S.D., CA 1985). The multiple cloning sites of pUC18 and pUC19 vectors were exchanged with those of pUC-CAM to create pCGN565 and pCGN566 which are CAM resistant. Also used were pUC118 and pUC119, which are respectively, pUC18 and pUC19 with the intergenic region of M13, from a HgiAl site at 5465 to the Ahalll site at 5941, inserted at the Ndel site of pUC. (Available from Vieira J. and Messing, J. Waksman Institute, Rutgers University, Rutgers, N.J.).

Materials

Terminal deoxynucleotide transferase (TDT), RNaseH, E. Coli, DNA polymerase, T4 kinase, and restriction enzymes were obtained from Bethesda Research Laboratories; E. coli DNA ligase was obtained from New England Biolabs; reverse transcriptase was obtained from Life Sciences, Inc.; isotopes were obtained from Amersham; X-gal was obtained from Bachem, Inc. Torrance, CA.

Construction of cDNA Library from Spinach Leaves

Total RNA was extracted from young spinach leaves in 4M guanidine thiocyanate buffer as described by Facciotti et al. (Biotechnology (1985) 3:241-246.). Total RNA was subjected to oligo(dT)-cellulose column chromatography two times to yield poly(A)+ RNA as described by Maniatis et al., Molecular Cloning: A Laboratory Manual. Cold Spring Harbor Laboratory, New York (1982). A cDNA library was constructed in pUC13-Cm according to the method of Gubler and Hoffman, (Gene (1983) 25:263-269) with slight modifications. RNasin was omitted in the synthesis of first strand cDNA as it interfered with second strand cDNA synthesis if not completely removed, and dCTP was used to tail the vector DNA and dGTP to tail

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The cDNA restriction, liga was prepared Nat. Acad. Sci IntelliGenetics

double-stranded cDNA instead of the reverse as described in the paper. The annealed cDNA was transformed to competent E. coli JM83 (Messing in Recombinant DNA Technical Bulletin, NIH Publication No. 79-99, 2 (1979) No.2:43-48.) cells according to Hanahan (J. Mol. Biol. (1983) 166:557-580) and spread onto LB agar plates (Miller, Experiments in Molecular Genetics (1972) Cold Spring Harbor Laboratory. Cold Spring Harbor, New York) containing 50µg/ml chloramphenicol and 0.005% X-Gal.

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Identification of Spinach ACP-I cDNA

A total of approximately 8000 cDNA clones were screened by performing Southern blots (Southern J. Mol. Biol (1975) 98:503) and dot blot (described below) hybridizations with clone analysis (see below) DNA from 40 pools representing 200 cDNA clones each. A 5' end-labeled synthetic oligonucleotide (ACPP4) that is at least 66% homologous with a 16 amino acid region of spinach ACP-I 5'-

CATGTCTTGAGCCTTGTCCTCATCCACATTGATACCAAACTCCTCCTC-3' is the complement to a DNA sequence that could encode the 16 amino acid peptide glu-glu-glu-phe-gly-ile-asn-val-asp-glu-asp-lys-ala-gln-asp-ile, residues 49-64 of spinach ACP-I (Kuo and Ohlrogge Arch. Biochem. Biophys. (1984) 234:290-296) and was used for an ACP proble.

Clone analysis DNA for Southern and dot blot hybridizations was prepared as follows. Transformants were transferred from agar plates to LB containing 50µg/ml chloramphenicol in groups of ten clones per 10/ml media. Cultures were incubated overnight in a 37° C shaking incubator and then diluted with an equal volume of media and allowed to grow for 5 more hours. Pools of 200 cDNA clones each were obtained by mixing contents of 20 samples. DNA was extracted from these cells as described by Birnboim and Doly (Nucleic Acids Res. (1979) 7:1513-1523). DNA was purified to enable digestion with restriction enzymes by extractions with phenol and chloroform followed by ethanol precipitation. DNA was resuspended in sterile, distilled water and 1µg of each of the 40 pooled DNA samples was digested with EcoRI and HindIII and electrophoresed through 0.7% agarose gels. DNA was transferred to nitrocellulose filters following the blot hybridization technique of Southern.

ACPP4 was 5' end-labeled using γ -32P dATP and T4 kinase according to the manufacturer's specifications. Nitrocellulose filters from Southern blot transfer of clone analysis DNA were hybridized (24 hours, 42°C) and washed according to Berent et al. (BioTechniques (1985) 3:208-220). Dot blots of the same set of DNA pools were prepared by applying 1µg of each DNA pool to nylon membrane filters in 0.5M NaOH. These blots were hybridized with the probe for 24 hours at 42°C in 50% formamide/1% SDS/1 M NaCl, and washed at room temperature in 2X SSC/0.1% SDS (1X SSC = 0.15M NaCl; 0.015M Na citrate; SDS-sodium dodecylsulfate). DNA from the pool which was hybridized by the ACPP4 oligoprobe was transformed to JM83 cells and plated as above to yield individual transformants. Dot blots of these individual cDNA clones were prepared by applying DNA to nitrocellulose filters which were hybridized with the ACPP4 oligonucleotide probe and analyzed using the same conditions as for the Southern blots of pooled DNA samples.

Nucleotide Sequence Analysis

The positive clone, pCGN1SOL, was analyzed by digestion with restriction enzymes and the following partial map was obtained.

pUC13-Cm	-35 *	248	1-63	- 152	 ~200	-1	
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**polylinker with available restriction sites indicated

The cDNA clone was subcloned into pUC118 and pUC119 using standard laboratory techniques of restriction, ligation, transformation, and analysis (Maniatis et al., (1982) supra). Single-stranded DNA template was prepared and DNA sequence was determined using the Sanger dideoxy technique (Sanger et al., Proc. Nat. Acad. Sci. (1977) USA 74:5463-5467). Sequence analysis was performed using a software package from IntelliGenetics, Inc.

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pCGN1SOL contains an (approximately) 700 bp cDNA insert including a stretch of A residues at the 3' terminus which represents the poly(A) tail of the mRNA. An ATG codon at position 61 is presumed to encode the MET translation initiation codon. This codon is the start of a 411 nucleotide open reading frame, of which, nucleotides 229-471 encode a protein whose amino acid sequence corresponds almost perfectly with the published amino acid sequence of ACP-I of Ohlrogge and Kuo <u>supra</u> sequence as described previously. Discrepancies between the two amino acid sequences are indicated in the sequence set forth previously. In addition to mature protein, the pCGN1SOL also encodes a 56 residue transit peptide sequence, as might be expected for a nuclear-encoded chloroplast protein.

Construction of a Napin Promoter

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There are 298 nucleotides upstream of the ATG start codon of the napin gene on the pgN1 clone (a 3.3 kb EcoRl fragment of B. napus genomic DNA containing a napin gene cloned into pUC8, available from Marti Crouch, University of Indiana), pgN1 DNA was digested with EcoRI and SstI and ligated to EcoRI/SstI digested pCGN706. (pCGN706 is an Xhol/Pstl fragment containing 3' and polyadenylation sequences of another napin cDNA clone pN2 (Crouch et al., 1983 supra) cloned in pCGN566 at the Sall and Pstl sites.) The resulting clone pCGN707 was digested with Sall and treated with the enzyme Bal31 to remove some of the coding region of the napin gene. The resulting resected DNA was digested with Smal after the Bal31 treatment and religated. One of the clones pCGN713, selected by size, was subcloned by EcoRi and BamHI digestion into both EcoRI/BamHI digested pEMBL18 (Dente et al., Nucleic Acids Res. (1983) 11:1645-1655) and pUC118 to give E418 and E4118 respectively. The extent of Bal31 digestion was confirmed by Sanger dideoxy sequencing of E418 template. The Bal31 deletion of the promoter region extended only to 57 nucleotides downstream of the start codon, thus containing the 5' end of the napin coding sequence and about 300 bp of the 5' non-coding region, E4118 was tailored to delete all of the coding region of napin including the ATG start codon by in vitro mutagenesis by the method of Zoller and Smith (Nucleic Acids Res. (1982) 10:6487-6500) using an oligonucleotide primer 5'-GATGTTTTGTATGTGGGCCCCTAGGAGATG-3'. Screening for the appropriate mutant was done by two transformations into E. coli strain JM83 (Messing, J., supra) and Smal digestion of putative transformants. The resulting napin promoter clone is pCGN778 and contains 298 nucleotides from the EcoRI site of pgN1 to the A nucleotide just before the ATG start codon of napin. The promoter region was subcloned into a chloramphenicol resistant background by digestion with EcoRI and BamHI and ligation to EcoRI/BamHI digested pCGN565 to give pCGN779c.

Extension of the Napin Promoter Clone

pCGN779c contains only 298 nucleotides of potential 5'-regulatory sequence. The napin promoter was extended with 1.8 kb fragment found upstream of the 5'-EcoRl site on the original λBnNa clone. The ~3.5 kb Xhol fragment of λBnNa (available from M. Crouch), which includes the napin region, was subcloned into Sall-digested pUC119 to give pCGN930. A HindIII site close to a 5' Xhol site was used to subclone the HindIII/EcoRl fragment of pCGN930 into HindIII/EcoRl-digested Bluescript + (Vector Cloning Systems, San Diego, CA) to give pCGN942. An extended napin promoter was made by ligating pCGN779c digested with EcoRl and PstI and pCGN942 digested with EcoRl and PstI to make pCGN943. This promoter contains ~2.1 kb of sequence upstream of the original ATG of the napin gene contained on λBnNa.

Napin Cassette

The extended napin promoter and a napin 3'-regulatory region is combined to make a napin cassette for expressing genes seed-specifically. The napin 3' region used is from the plasmid pCGN1924 containing the Xhol/EcoRl fragment from pgN1 (Xhol site is located 18 nucleotides from the stop codon of the napin gene) subcloned into EcoRl/Sall digested pCGN565. HindIll/Pstl digested pCGN943 and pCGN1924 are ligated to make the napin cassette pCGN944, with unique cloning sites Smal, Sall and Pstl for inserting genes.

Napin - ACP Construct

pCGN796 was constructed by ligating pCGN1SOL digested with HindIII/BamHI. pUC8 digested with HindIII and BamHI and pUC118 digested with BamHI. The ACP gene from pCGN796 was transferred into a chloramphenicol background by digestion with BamHI and ligation with BamHI digested pCGN565. The resulting pCGN1902 was digested with EcoRI and Smal and ligated to EcoRI/Smal digested pUC118 to give pCGN1920. The ACP gene in pCGN1920 was digested at the NcoI site, filled in by treatment with the Klenow fragment, digested with Smal and religated to form pCGN1919. This eliminated the 5'-coding sequences from the ACP gene and regenerated the ATG. This ACP gene was flanked with PstI sites by digesting pCGN1919 with EcoRI, filling in the site with the Klenow fragment and ligating a PstI linker. This clone is called pCGN945. The ACP gene of pCCGN945 was moved as a BamHI/PstI fragment to pUC118 digested with BamHI and PstI to create pCGN945a so that a Smal site (provided by the pUC118) would be at the 5'-end of the ACP sequence to facilitate cloning into the napin cassette pCGN944. pCGN945a digested with Smal and PstI was ligated to pCGN944 digested with Smal and PstI to produce the napin ACP cassette pCGN946. The napin ACP cassette was then transferred into the binary vector pCGN783 by cloning from the HindIII site to produce pCGN948.

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pCGN783 is a binary plasmid containing the left and right T-DNA borders of A. tumefaciens (Barker et al. Plant Mol. Biol. (1983) 2:335-350); the gentamicin resistance gene of pPH1JI (Hirsch et al., (1984) Plasmid 12. 139-141) the 35S promoter of cauliflower mosaic virus (CaMV) (Gardner et al., Nucleic Acids Res. (1981) 9:2871-2890), the kanamycin resistance gene of Tn5 (Jorgensen et al., infra and Wolff et al. ibid (1985) 13:355-367) and the 3' region from transcript 7 of pTiA6 (Barker et al., (1983) supra).

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To obtain the gentamicin resistance marker, the gentamicin resistance gene was isolated from a 3.1 kb EcoRI-PstI fragment of pPHIJ1 and cloned into pUC9 yielding pCGN549. The HindIII-BamHI fragment containing the gentamicin resistance gene was substituted for the HindIII-BgIII fragment of pCGN587 creating pCGN594.

pCGN587 was prepared as follows: The HindIII-Smal fragment of Tn5 containing the entire structural gene for APHII (Jorgensen et al., Mol. gen. Genet. (1979) 177:65) was cloned into pUC8 (Vieira and Messing, Gene (1982) 19:259), converting the fragment into a HindIII-EcoRI fragment, since there is an EcoRI site immediately adjacent to the Smal site. The PstI-EcoRI fragment containing the 3'-portion of the APHII gene was then combined with an EcoRI-BamHI-Sall-PstI linker into the EcoRi site of pUC7 (pCGN546W). Since this construct does not confer kanamycin resistance, kanamycin resistance was obtained by inserting the Bg1II-pstI fragment of the APHII gene into the BamHI-PstI site (pCGN546X). This procedure reassembles the APHII gene, so that EcoRI sites flank the gene. An ATG codon was upstream from and out of reading frame with the ATG initiation codon of APHII. The undesired ATG was avoided by inserting a Sau3A-PstI fragment from the 5'-end of APHII, which fragment lacks the superfluous ATG, into the BamHI-PstI site of pCGN546W to provide plasmid pCGN550.

The EcoRI fragment containing the APHII gene was then cloned into the unique EcoRI site of pCGN451, which contains an octopine synthase cassette for expression, to provide pCGN552 (1ATG).

pCGN451 includes an octopine cassette which contains about 1556 bp of the 5' non-coding region fused via an EcoRI linker to the 3' non-coding region of the octopine synthase gene of pTiA6. The pTi coordinates are 11,207 to 12,823 for the 3' region and 13,643 to 15,208 for the 5' region as defined by Barker et al., Plant Mol. Biol. (1983) 2:325.

The 5' fragment was obtained as follows. A small subcloned fragment containing the 5' end of the coding region, as a BamHI-EcoRI fragment was cloned in pBR322 as plasmid pCGN407. The BamHI-EcoRi fragment has an XmnI site in the coding region, while pBR322 has two XmnI sites. pCGN407 was digested with Xmnf, resected with Ba131 nuclease and EcoRI linkers added to the fragments. After EcoRI and BamHI digestion, the fragments were size fractionated, the fractions cloned and sequenced. In one case, the entire coding region and 10 bp of the 5' non-translated sequences had been removed leaving the 5' non-transcribed region, the mRNA cap site and 16 bp ofthe 5' non-translated region (to a BamHI site) intact. This small fragment was obtained by size fractionation on a 7% acrylamide gel and fragments approximately 130 bp long eluted.

This size fractionated DNA was ligated into M13mp9 and several clones sequenced and the sequence compared to the known sequence of the octopine synthase gene. The M13 construct was designated p14, which plasmid was digested with BamHI and EcoRI to provide the small fragment which was ligated to a Xhol to BamHI fragment containing upstream 5' sequences from pTiA6 (Garfinkel and Nester, J. Bacteriol. (1980) 144:732) and to an EcoRI to Xhol fragment containing the 3' sequences.

The resulting Xhol fragment was cloned into the Xhol site of a pUC8 derivative, designated pCGN426. This plasmid differs from pUC8 by having the sole EcoRI site filled in with DNA polymerase I, and having lost the PstI and HindIII site by nuclease contamination of HincII restriction endonuclease, when a Xhol linker was inserted into the unique HincII site of pUC8. The resulting plasmid pCGN451 has a single EcoRI site for the insertion of protein coding sequences between the 5' non-coding region (which contains 1,550 bp of 5' non-transcribed sequence including the right border of the T-DNA, the mRNA cap site and 16 bp of 5' non-translated sequence) and the 3' region (which contains 267 bp of the coding region, the stop codon, 196 bp of 3' non-translated DNA, the polyA site and 1,153 bp of 3' non-transcribed sequence). pCGN451 also provides the right T-DNA border.

The resulting plasmid pCGN451 having the ocs 5' and the ocs 3' in the proper orientation was digested with EcoRI and the EcoRI fragment from pCGN551 containing the intact kanamycin resistance gene inserted into the EcoRI site to provide pCGN552 having the kanamycin resistance gene in the proper orientation.

This ocs/KAN gene was used to provide a selectable marker for the trans type binary vector pCGN587. The 5' portion of the engineering octopine synthase promoter cassette consists of pTiA6 DNA from the Xhol at bp 15208-13644 (Barker's numbering), which also contains the T-DNA boundary sequence (border) implicated in T-DNA transfer. In the plasmid pCGN587, the ocs/KAN gene from pCGN552 provides a selectable marker as well as the right border. The left boundary region was first cloned in M13mp9 as a HindIII-Smal piece (pCGN502) (base pairs 602-2213) and recloned as a KpnI-EcoRI fragment in pCGN565 to provide pCGN580. pCGN565 is a cloning vector based on pUC8-Cm, but containing pUC18 linkers. pCGN580 was linearized with BamHI and used to replace the smaller Bg1II fragment of pVCK102 (Knauf and Nester, Plasmid (1982) 8:45), creating pCGN585. By replacing the smaller Sa1I fragment of pCGN585 with the Xhol fragment from pCGN552 containing the ocs/KAN gene, pCGN587 was obtained.

The pCGN594 <u>HindIII-BamHI</u> region, which contains an 5'-ocs-kanamycin-ocs-3' (ocs is octopine synthase with 5' designating the promoter region and 3' the terminator region, see U.S. application serial no. 775,923, filed September 13, 1985) fragment was replaced with the HindIII-BamHI polylinker region from pUC18.

pCGN566 contains the EcoRI to HindIII polylinker of pUC18 inserted into the EcoRI-HindIII sites of pUC13-Cm. The HindIII-BgIII fragment of pNW31C-8,29-1 (Thomashow et al., Cell (1980) 19:729) containing ORF1 and -2 of pTiA6 was subcloned into the HindIII-BamHI sites of pCGN566 producing pCGN703.

The Sau3A fragment of pCGN703 containing the 3' region of transcript 7 (corresponding to bases 2396-2920 of pTiA6 (Barker et al., (1983) supra) was sub-cloned into the BamHI site of pUC18 producing pCGN709. The EcoRI-Smal polylinker region of pCGN709 was substituted with the EcoRI-Smal fragment of pCGN587, which contains the kanamycin resistance gene (APH3-II) producing pCGN726.

The EcoRI-Sa1I fragment of pCGN726 plus the BgIII-EcoRI fragment of pCGN734 were inserted into the BamHI-SaII site of pUC8-Cm producing pCGN738. pCGN726c is derived from pCGN738 by deleting the 900 bp EcoRI-EcoRI fragment.

To construct pCGN167, the Alul fragment of CaMV (bp 7144-7735) (Gardner et al., (1981) supra) was obtained by digestion with Alul and cloned into the Hincll site of M13mp7 (Messing, et al., Nucl. Acid Res. (1981) 9:309-321) to create C614. An EcoRl digest of C614 produced the EcoRl fragment from C614 containing the 35S promoter which was cloned into the EcoRl site of pUC8 (Vieira and Messing et al., Gene (1982) 19:259) to produce pCGN146.

To trim the promoter region, the <u>Bglll</u> site (bp 7670) was treated with <u>Bglll</u> and resected with <u>Bal31</u> and subsequently a <u>Bglll</u> linker was attached to the <u>Bal31</u> treated DNA to produce pCGN147.

pCGN148a containing a promoter region, selectable marker (KAN with 2 ATG's) and 3' region, was prepared by digesting pCGN528 with Bglll and inserting the BamHI-Bglll promoter fragment from pCGN147. This fragment was cloned into the Bglll site of pCGN528 so that the Bglll site was proximal to the kanamycin gene of pCGN528.

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The shuttle vector used for this construct, pCGN528, was made as follows, pCGN525 was made by digesting a plasmid containing Tn5 which harbors a kanamycin gene (Jorgenson et al., Mol. gen. Genet. (1979) 177:65) with HindIII-BamHI and inserting the HindIII-BamHI fragment containing the kanamycin gene into the HindIII-BamHI sites in the tetracycline gene of pACYC184 (Chang & Cohen, J. Bacteriol. (1978) 134:1141-1156), pCGN526 was made by inserting the BamHI fragment 19 of pTiA6 (Thomashow et al., Cell (1980) 19:729-739), modified with Xhol linkers inserted into the Smal site, into the BamHI site of pCGN525, pCGN528 was obtained by deleting the small Xhol fragment from pCGN526 by digesting with Xhol and religating.

pCGN149a was made by cloning the <u>Bam</u>HI-kanamycin gene fragment from pMB9KanXXI into the <u>Bam</u>HI site of pCGN148a.

pMB9KanXXI is a pUC4K variant (Vieira and Messing, Gene (1982) 19:259-268) which has the XhoI site missing but contains a functional kanamycin gene from Tn903 to allow for efficient selection in Agrobacterium.

pCGN149a was digested with Bglll and Sphl. This small Bglll-Sphl fragment of pCGN149a was replaced with the BamHI-Sphl fragment from MI (see below) isolated by digestion with BamHI and Sphl. This produces pCGN167, a construct containing a full length CaMV promoter, 1ATG-kanamycin gene, 3' end and the bacterial Tn903-type kanamycin gene. MI is an EcoRI fragment from pCGN546X (see construction of pCGN587) and was cloned into the EcoRI cloning site of M13 mp9 in such a way that the Pstl site in the 1ATG-kanamycin gene was proximal to the polylinker region of M13mp9.

The HindIII-BamHI fragment in the pCGN167 containing the CaMV-35S promoter, 1ATG-kanamycin gene and the BamHI-fragment 19 of pTiA6 was cloned into the BamHI-HindIII sites of pUC19 creating pCGN976. The 35S promoter and 3' region from transcript 7 was developed by inserting a 0.7 kb HindIII-EcoRI fragment of pCGN976 (35S promoter) and the 0.5 kb EcoRI-Saul fragment of pCGN709 (transcript 7:3') into the HindIII-Sall sites of pCGN566 creating pCGN766c.

The 0.7 kb HindIII-EcoRI fragment of pCGN766c (CaMV-35S promoter) was ligated to the 1.5 kb EcoRI-Sall in pCGN726c (1ATG-KAN 3' region) into the HindIII-Sall sites of pUC119 to produce pCGN778. The 2.2 kb region of pCGN778, HindIII-Sall fragment containing the CaMV-35S promoter and 1ATG-KAN-3' region was used to replace the HindIII-Sall linker region of pCGN739 to produce pCGN783.

pCGN948 was introduced into Agrobacterium tumefaciens EHA101 (Hood et al, J. Bacteriol. (1986) 168:1291-1301) by transformation. An overnight two ml culture of EHA101 was grown in MG/L broth at 30°C. 0.5 ml was inoculated into 100 ml of MG/L broth (Garfinkel and Nester, J. Bacteriol. (1980) 144:732-743) and grown in a shaking incubator for 5 h at 30°C. The cells were pelleted by centrifugation at 7K, resuspended in 1 ml of MG/L broth and placed on ice. Approximately one µg of pCGN948 DNA was placed in 100 µl of MG/L broth to which 200 µl of the EHA101 suspension was added; the tube containing the DNA-cell mix was immediately placed into a dry ice/ethanol bath for 5 minutes. The tube was quick thawed by 5 minutes in a 37°C water bath followed by 2h of shaking at 30°C after adding 1 ml of fresh MG/L medium. The cells were pelleted and spread onto MG/L plates (1.5% agar) containing 100 mg/l gentamicin. Plasmid DNA was isolated from individual genta micin-resistant colonies, transformed back into E.coli, and characterized by restriction enzyme analysis to verify that the gentamicin-resistant EHA101 contained intact copies of pCGN948. Single colonies are picked and purified by two more streakings on MG/L plates containing 100 mg/l gentamicin.

Seeds of <u>Brassica napus</u> of Westar were soaked in 95% ethanol for 4 minutes. They were sterilized in 1% solution of sodium hypochlorite with 50 μl of "Tween 20" surfactant per 100 ml sterilent solution. After soaking for 45 minutes, seeds were rinsed 4 times with sterile distilled water. They were planted in sterile plastic boxes 7 cm wide, 7 cm long, and 10 cm high (Magenta) containing 50 ml of 10/10th concentration of MS (Murashige minimal organics medium, Gibco) with added pyridoxine (50 μg/l), nicotinic acid (50 μg/l), glycine

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(200 µg/I) and solidified with 0.6% agar. The seeds germinated and were grown at 22°C in a 16h-8h light-dark sites of cycle with light intensity approximately 65 µEm-2s-1. After 5 days, the seedlings were taken under sterile ntaining conditions and the hypocotyls excised and cut into pieces of about 4 mm in length. The hypocotyl segments `03. were placed on a feeder plat or without the feeder layer on top of a filter paper on the solidified B5 0/1/1 or 36-2920 B5 0/1/0 medium. B5 0/1/0 medium contains B5 salts and vitamins (Gamborg, Miller and Ojima, Experimental '09. The Cell Res. (1968) 50:151-158), 3% sucrose. 2,4-dilchlorophenoxyacetic acid (1.0 mg/l), pH adjusted to 5.8, and , which the medium is solidified with 0.6% Phytagar; B5 0/1/1 is the same with the addition of 1.0 mg/l kinetin. Feeder plates were prepared 24 hours in advance by pipetting 1.0 ml of a stationary phase tobacco suspension culture into the (maintained as described in Fillatti et al., Mol. gen. Genet. (1987) 206:192-199) onto B5 0/1/0 or B5 0/1/1 medium. Hypocotyl segments were cut and placed on feeder plates 24 hours prior to Agrobacterium

> Agrobacterium tumefaciens (strain EHA101 x 948) were prepared by incubating a single colony of Agrobacterium in MG/L broth at 30°C. Bacteria were harvested 16 hours later and dilutions of 108 bacteria per ml were prepared in MG/L broth. Hypocotyl segments were inoculated with bacteria by placing in Agrobacterium suspension and allowed to sit for 30-60 minutes, then removed and transferred to Petri plates containing B5 0/1/1 or 0/1/0 medium described above. The plates were incubated in low light at 22°C. The co-incubation of bacteria with the hypocotyl segments took place for 24-48 hours. The hypocotyl segments were removed and placed on B5 0/1/1 or 0/1/0 containing 500 mg/1 carbenicillin (kanamycin sulfate at 10, 25. or 50 mg/l was sometimes added at this time) for 7 days in continuous light (approximately 65 μ Em-2S-1) at 22°C. They were transferred to B5 salts medium containing 1% sucrose, 3 mg/1 benzylamino-purine and 1 mg/1 zeatin. This was supplemented with 500 mg/1 carbenicillin, 10, 25, or 50 mg/1 kanamycin sulfate, and solidified with 0.6% Phytagar (Gibco). Thereafter explants were transferred to fresh medium every 2 weeks.

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After 1 month green shoots developed from green calli which were selected on media containing kanamycin. Shoots continued to develop for 3 months. The shoots were cut from the calli when they were at least 1 cm high and placed on B5 medium with 1% sucrose, no added growth substances, 300 mg/1 carbenicillin, and solidified with 0.6% phytagar. The shoots continued to grow and several leaves were removed to test for neomycin phosphotransferase II (NPTII) activity. Shoots which were positive for NPTII activity were placed in Magenta boxes containing B5 0/1/1 medium with 1% sucrose, 2 mg/1 indolebutyric acid, 200 mg/1 carbenicillin, and solidified with 0.6% Phytagar. After a few weeks the shoots developed roots and were transferred to soil. The plants were grown in a growth chamber at 22°C in a 16-8 hours light-dark cycle with light intensity 220 μEm-2s-1 and after several weeks were transferred to the greenhouse.

SOUTHERN DATA

treatment.

Regenerated B. napus plants from cocultivations of Agrobacterium tumefaciens EHA101 containing pCGN948 and B. napus hypocotyls were examined for proper integration and embryo-specific expression of the spinach leaf ACP gene. Southern analysis was performed using DNA isolated from leaves of regenerated plants by the method of Dellaporta et al. (Plant Mol. Biol. Rep. (1983) 1:19-21) and purified once by banding in CsCl. DNA (10 µg) was digested with the restriction enzyme EcoR1, electrophoresed on a 0.7% agarose gel and blotted to nitrocellulose (see Maniatis et al., 1982, supra). Blots were probed with pCGN945 DNA containing 1.8 kb of the spinach ACP sequence or with the EcoRI/HindIII fragment isolated from pCGN936c (made by transferring the HindIII/EcoRI fragment of pCGN930 into pCGN566) containing the napin 5' sequences labeled with 32P-dCTP by nick translation (described by the manufacturer, BRL Nick Translation Reagent Kit, Bethesda Research Laboratories, Gaithersburg, MD.). Blots were prehybridized and hybridized in 50% formamide, 10xDenhardt's, 5xSSC, 0.1% SDS, 5mM EDTA, 100µg/ml calf thymus DNA and d10% dextran sulfate (hybridization only) at 42°C. (Reagents described in Maniatis et al., (1982) supra). Washes were in 1xSSC, 0.1% SDS, 30 min and twice in 0.1xSSC, 0.1% SDS at 55°C.

Autoradiograms showed two bands of approximately 3.3 and 3.2 kb hybridized in the EcoRI digests of DNA from four plants when probed with the ACP gene (pCGN945) indicating proper integration of the spinach leaf ACP construct in the plant genome since 3.3 and 3.2 kb EcoRI fragments are present in the T-DNA region of pCGN948. The gene construct was present in single or multiple loci in the different plants as judged by the number of plant DNA-construct DNA border fragments detected when probed with the napin 5' sequences.

NORTHERN DATA

Expression of the integrated spinach leaf ACP gene from the napin promoter was detected by Northern analysis in seeds but not leaves of one of the transformed plants shown to contain the construct DNA. Developing seeds were collected from the transformed plant 21 days post-anthesis. Embryos were dissected from the seeds and frozen in liquid nitrogen. Total RNA was isolated from the seed embryos and from leaves of the transformed plant by the method of Crouch et al. (1983) supra), electrophoresed on formaldehyde-containing 1.5% agarose gels as described (Shewmaker et al., Virology (1985) 140:281-288) and blotted to nitro-cellulose (Thomas, Proc. Natl. Acad. Sci., U.S.A. (1980) 77:5201-5205). Blots were prehybridized. hybridized and washed as described above. The probe was an isolated Pstl/BamHI fragment from pCGN945 containing only spinach leaf ACP sequences labeled by nick translation.

An RNA band of ~0.8 kb was detected in embryos but not leaves of the transformed plant indicating seed-specific expression of the spinach leaf ACP gene.

Although higher plant fatty acid biosynthetic genes have been shown to recognize acyl-ACP substrates

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when the ACP moiety is an Escherichia coli ACP, it remains possible that different forms of ACP from different sources may affect the efficiency and/or the final products of fatty acid synthesis in vivo. For instance, both spinach (Ohlrogge and Kuo, . Biol. Chem. (1985) 260:8032-8037) and barley (Hoj and Svendsen, Carlsberg. Res. Commun. (1984) 49:483-492, contain more than one form of ACP. To support the generality of using isoforms of ACP to enhance the value of oilseed crops, a cDNA copy of the gene for an ACP found in the seeds of the oilseed crop turnip rape (Brassica campestris) was isolated and characterized.

The following is the procedure for integrating the ACP gene into a chimeric gene using a napin promoter from Brassica campestris rather than Brassica napus.

Immature seeds are collected from Brassica campestris cv. "R-500", a self-compatible variety of turnip rape. Whole seeds are collected at stages corresponding approximately to 14 to 28 days after flowering. RNA isolation and preparation of a cDNA bank was as described above for the isolation of a spinach ACP cDNA clone. To probe the cDNA bank, the oligonucleotide (5')-ACTTTCTCAACTGTCTCTGGTTTAGCAGC-(3) was synthesized using an Applied Biosystems DNA Synthesizer, model 380A, according to manufacturer's recommendations. This synthetic DNA molecule will hybridize at low stringencies to DNA or RNA sequences coding for the amino acid sequence (ala-ala-lys-pro-glu-thr-val-glu-lys-val). This amino acid sequence has been reported for ACP isolated from the seeds of Brassica napus (Slabas et al., 7th International Symposium of the Structure and Function of Plant Lipids, University of California, Davis, CA, Plenum Press, N.Y. 1987); the ACP from B. campestrisseed is highly homologous. Approximately 2200 different cDNA clones are analyzed using a colony hybridization technique (Taub and Thompson, Anal. Biochem. (1982) 126:222-230) and hybridization conditions corresponding to (Wood et al., Proc. Natl. Acad. Sci. U.S.A. (1985) 82:1585-1588). DNA sequence analysis of two cDNA clones showing obvious hybridization to the oligonucleotide probe indicated that one, designated pCGN1Bcs, coded for an ACP-precursor protein as evidenced by the considerable homology of the encoded amino acid sequence with ACP proteins described from Brassica napus (Slabas et al., supra). The DNA sequence of pCGN1Bcs (referred to also as AGB1) is indicated as follows:

AGB1 CD

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70 CCACT

139 CCTTC erPhe

208 AGAAA luLys

277 AGTTT ysPhe

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	218 237	
	Sau3AI HinfI TagI	40
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•	277 ACMMMCCMCACCACCACCAMMCMCCACCACCACCACCACC	
	277 AGTTTGCTGATCTTGGAGCAGATTCTCTCGACACTGTTGAGATAGTGATGGGTTTAGAGGAAGAGTTTG 345	
	ysPheAlaAspLeuGlyAlaAspSerLeuAspThrValGluIleValMETGlyLeuGluGluGluPheA	
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	346 ATATCGAAATGGCTGAAGAGAAGCTCAGAAGATTGCTACTGTGGAGGAAGCTGCTGAACTCATTGAAG 414	
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	348 370 397	
	350 371	
		55

SacI AluI 1 1 415 AGCTCGTTCAACTTAAGAAGTAATTTTAGTATTAAGAGCAGCCAAGGCTTTGTTGGGTTTTGTTTTC 483 5 luLeuValGlnLeuLysLys . 417 419 10 HinfI DraI 484 ATAATCTTCCTGTCATTTTCTTTTTCTTTAATGTGTCAAGCGACTCTGTTGGTTTAAAGTAGTATCTGT 552 526 539 15 553 TTGCCAAAAAA 564 To achieve high-level embryo-specific expression of a Brassica campestris seed ACP in a transgenic 20 Brassica napus, a chimeric gene is made analogous to the embryo-specific chimeric gene employing spinach ACP-coding DNA sequences as described for pCGN946 above. The pCGN1Bcs ACP-coding region is adapted to fit into the Brassica campestris napin-type promoter element present in pCGN1803 (described below). CONSTRUCTION OF B. CAMPESTRIS NAPIN PROMOTER CASSETTE 25 A Bglll-partial genomic library of B. campestris DNA was made in the lambda vector Charon 35 using established protocols (Miniatis et al., (1982) supra). The titer of the amplified library was ~1.2x109 phage/ml. Four hundred thousand recombinant bactriophage were plated at a density of 105 per 9x9 NZY plate (NZYM as described in Maniatis et al., 1982 supra) in NZY + 10mM MgSO₄ + 0.9% agarose after adsorption to DH1 E. coli cells (Hanahan, D., J. Mol. Biol. (1983) 166:557) for 20 min at 37°C. Plates were incubated at 37°C for -13 hours, cooled at 4°C for two and one half hours and phage were lifted onto GeneScreen Plus (New England Nuclear) by laying precut filters over the plates for approximately 1 min and peeling them off. The adsorbed phage DNA was immobilized by floating the filter on 1.5M NaCl, 0.5M NaOH for 1 min., neutralizing in 1.5M NaCl, 0.5M Tris-HCl, pH 8.0 for 2 min and 2XSSC for 3 min. Filters were air dried until just damp, prehybridized and hybridized at 42°C as described for Southern analysis. Filters were probed for napin-containing clones 35 using an Xhol/Sall fragment of the cDNA clone BE5 which was isolated from the B. campestris seed cDNA library described above using the probe pN1 (Crouch et al., 1983, supra). Three plaques were hybridized strongly on duplicate filters and were plaque purified as described (Maniatis et al., 1982 supra). The following is the BE5 cDNA sequence. 40 45 50

BES CDN

1 ACAGA ThrG1

70 ACCAAT

ThrAsc

139 TTTAGG

208 CACAAG(HisLys(

277 GACGTGG AspValG

Bst | |346 CACCAGG |HisGlnG

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BE5 CDNA LENGTH = 734 AluI 1 ACAGAACATACACAAATGGCGAACAAGCTCTTCCTCGTCTCGGCCAACTCTCGCCTTGTTCTTCTTCTC ThrGluHisThrGlnMETAlaAsnLysLeuPheLeuValSerAlaThrLeuAlaLeuPhePheLeuLeu 28 TaqI SalI NaeI HincII Hpall ACCT AccI HaeIII 70 ACCANTGCCTCCGTCTACAGGACGGTTGTGGAAGTCGACGAAGATGCCACAAATCCAGCCGGCCCA 138 ThrasnalaSerValTyrArgThrValValGluValAspGluAspAspAlaThrAsnProAlaGlyPro 84 105 103 132 104 133 105 .genic 20 ∋inach apted HindIII elow). AluI 139 TTTAGGATTCCAAAATGTAGGAAGGAGTTTCAGCAAGCACAACACCTGAAAGCTTGCCAACAATGGCTC 207 PheArgIleProLysCysArgLysGluPheGlnGlnAlaGlnHisLeuLysAlaCysGlnGlnTrpLeu 25 usina 191 ge/ml. 189 YM as)H1 E. Avall or ~13 HpaII 30 AvaII TaqI AluI igland 1 208 CACAAGCAGGCAATGCAGTCCGGTAGTGGTCCAAGCTGGACCCTCGATGGTGAGTTTGATTTTGAAGAC 276 orbed HisLysGlnAlaMETGlnSerGlySerGlyProSerTrpThrLeuAspGlyGluPheAspPheGluAsp 1.5M ridized 228 236 243 252 clones 35 cDNA ridized SacI HaeIII HaeIII AluI 277 GACGTGGAGAACCAACAACAGGGCCCGCAGCAGGGCCCACCGCTGCTCCAGCAGTGCTGCAACGAGCTC 345 AspValGluAsnGlnGlnGlnGlyProGlnGlnArgProProLeuLeuGlnGlnCysCysAsnGluLeu 40 300 313 343 345 **BstNI** 346 CACCAGGAAGAGCCACTTTGCGCTTTGCCCAACCTTGAAAGGAGCATCCAAAGCCGTTAAACAACAGATT 414 HisGlnGluGluProLeuCysValCysProThrLeuLysGlyAlaSerLysAlaValLysGlnGlnIle 45

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5		415		• <i>:</i>	x
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10	404	AlaThrHisLeuProArgAlaCysAsnIl 503	eArgGlnValSerI	eCysProPheGlnLysThrM2TPro 55	
15		Hpall HaellI GGGCCCGGCTTCTACTAGATTCCAAACGA	XhoI TaqI AvaI 	AccI 	. 2
20	553	GlyProGlyPheTyr 556 558	588 589 588	600	70 AC
	622	HidclI GTTGTTGATGTATGTTAACACTACATAGT	CATGGTGTGTGTT	RsaI CATAAATAATGTACTAATGTAATAAG	690 139 TT
25		635		677	
30	601	AccI - AACTACTCCGTAGACGGTAATAAAAGAGA	•	raI	208 TG
	931			33	
<i>35</i>		702		33	277 TA
40					346 AA
45	OVE	One of the clones named lambda CGN1 erlapping 2.7 kb Xhol and 2.1 kb <u>Sal</u> l restr N1-2 DNA into pCGN789 (a pUC based v	iction fragments. T	e two fragments were subcloned fro	m lambda

GGAATTCGTCGACAGATCTCTGCAGCTCGAGGGATCCAAGCTT 3'-which represents the polylinker EcoRI. Sall, Bglll, Pstl, Xhol, BamHl, Hindlll). The identity of the subclones as napin was confirmed by sequencing. The entire coding region sequence as well as extensive 5' upstream and 3' downstream sequences were determined.

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Lambda CGN1-2		
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208 TGTAGGTTGGGCAAAAACGAGGAAGATTGCTTCTCAATTTGGAAGAAGAACAGCCGAAGAAGAAAA	276	30
Sau3AI DdeI 277 TAAGAATAGGCAGTCCTGCTACTCAATGGATCTCAGTCTATAACGGTCGTCGCCATGAAACAGAGGT	345	<i>35</i>
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EcoRV 346 AAAACATTTTTTGCATATACACTTTGAAAGTTCCTCACTAACTGTGTAATCTTTTGGTAGATATCACTA	414	40
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25	553 TGAGTTGTCACCGGTCTTCCTACACAAGGTAATAATC 564 564	AGTTGAAGCAATTAAGAATCAATTTGATTTGT 621 606	
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35	629 Alui Alui	Saci Alui AGCTCTTTAGTTGTTATGTCAAAAGGTTAGTGT 759	1243 CTCA
40	702 710	729 731	,
45	760 TTAGTGAATAATAAACTTATACCACAAAGTCTTCAT	TGACTTATTTATATACTTGTTGTGAATTGCTAG 828	1312 TAGG.
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	1548	
	n.abu	2140 G;
15	AluI EcoRV	
	1588 TTTGGGAGCTTTTTAAGCCCTTCAAGTGTGCTTTTTATCTTATTGATATCATTCAT	
	1596 1635	
20	Xbal DdeI	
	1657 TGCGTCTCTAGATATGTTCCTATATCTTTCTCAGTGTCTGATAAGTGAAAATGTGAGAAAACCATACCAA 1725	2209 AJ AE
25	1664 1687	
	HinfI	
30	1761	
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35	1795 TGCTGAATCTATCACACTAGAAAAAACATTTCTTCAAGGTAATGACTTGTGGACTATGTTCTGAATTC 1863	
	1800 1859	
40	1864 TCATTAAGTTTTTATTTTCTGAAGTTTAAGTTTTTACCTTCTGTTTTGAAATATATCGTTCATAAGATG 1932	2347 AA Ly
	SphI	•
45	BatNI AluI Sau3AI	
70	1933 TCACGCCÁGGACATGAGCTACACATCGCACATAGCATGCAGATCAGGACGATTTGTCACTCAC	
	1940 1950 1973 1971	

		DdeI	AluI	HhaI	NdeI	SphI NeiI	Sau3AI	
518	200:	2 CACCTAAGA	GCTTCTCTCTCAC	CAGCGCACACI	 ACATATGC	 ATGCAATATTT	l ACACGIGATCGCCATGCAA	207C
		2006	2012	2028	2036	2042	2058	5
	207:	1 ATCTCCATT	CTCACCTATAAA1	TAGAGCCTCS	GCTTCAC	TCTTTACTCAN	ACCAAAACTCATCACTACA	2139
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	2140) GAACATACA	CAAATGGCGAACA METAlaAsnL	 AGCTCTTCCT	CGTCTCG(uValSer)	GCAACTCTCGCC AlaThrLeuAls	TTGTTCTTCCTTCTCACC	2208
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	2278	AGGATTCCAP ArgileProf 2281	AATGTAGGAAGG LyaCyaArglyaG	AGTTTCAGCA luPheGlnGl	AGCACAAC nAlaGlnH	TEPROTIANTE	TGCCAACAATGGCTCCAC CyeGlnGlnTrpLeuHis	2346
963						2325		35
932			MapI Hpall	AvaII ;	1 1	TaqI		
332	2347	AAGCAGGCAA LysGlnAlaM	TGCAGTCĊGGTA ŒTGlnSerGlyS 2364 2364	G + G T A L T C C E	CTGGACCC rTrpThrL 2379 2382	TCGATGGTGAG BuAspGlyGlu 2388	TTTGATTTTGAAGACGAC : PheAspPheGluAspAsp	2415 ₄₀
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			HaeIII ApaI 	HaeIII		S Alu I I	SacI II	289
5	2416	GTGGAGAACCAAC ValGluAsnGlnG	AACAGGGCCGGCAGC InglnglyProGlnG 2438 2436	AGAGGCCACCGCTG 1nArgProProLeu 2449	CTCCAGCAG LeuGlnGln	CysCysKanGluber 247	nia	-0,
10	2485	BstNI CAGGAAGAGCCAG GlnGluGluProI 2486	TTTGCGTTTGCCCAA LauCysValCysProT	.CCTTGAAAGGAGCA hrleulysGlyAls	ATCCAAAGCC ASGTLYBALB) GTTAAACAACAGAT	ewid	296;
15	2554	CAACAACAGGGA GinGlnGlnGly	CAACAAATGCAGGGAC GlnGlnMETGlnGlyG	LAGCAGATGCAGCAI SINGINMETGINGI	AGTGATTAGO nVallle5e:	CGTATCTACCAGAC ArgileTyrGlnTh	CGCT 2622	303-
20	2623	ACGCACTTACCT	AluI 	AGGCAAGTTAGCAT ArgGlnValSeril	TTGCCCCTT(eCysProPh(CAGAAGACCATGCC	BstNI I TGGG 2691 CGLy 2688	3106
25								
		MspI HpaII RaeIII ApaI	Hinfl	XhoI TaqI AvaI 	AccI			3175
30	2692	2 CCCGGCTTCTAC ProGlyPheTyr 2694 2692	TAGATTCCAAACGAA	TATOCTCGAGAGTG 2724 2725	etgtätacca 2736	CGGTGATATGAGTG	IGGTT 2760	3244
35		2694 2694	·	2724				
		•	incII			Raal 		3313
40	276	1 GTTGATGTATG	TTAACACTACATAGTO	LATGGTGTGTGTTC	CATAAATAAT	•	AGAAC 2829	
		2.	771			2813		
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45	283	O TACTCCGTAGA	CGGTAATAAAAGAGAJ	AGITITTTTTTIA	CTCTTGCTAC	TTTCCTATAAAGTG	ATGAT 2898	
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2899	FANCANCAGATACACCANNAGANNACANTRATCTATATTCACAATGAAGCAGTACTAGTCTATTGAA 296	57
	2954 2954	5
2968	Sau3AI CATGTCAGATTTTCTTTTTCTAAATGTCTAATTAAGCCTTCAAGGCTAGTGATGATAAAGATCATCCA 303 3028	36 ¹⁰
3037	Sau3AI Sau3AI BamHI HinfI BclI 7 ATGGGATCCAACAAAGACTCAAATCTGGTTTTGATCAGATACTTCAAAACTATTTTTGTATTCATTAAA 3:0	<i>15</i> 05
	3041 3053 3069 3041 3069	20
3106	Hinfi TTATGCAAGTGTTCTTTTATTTGGTGAAGACTCTTTAGAAGCAAGC	7 4 25
	3135	
3175	S ACARAGITCAGITITAAGAITIGITATIGACITATIGICATITGAAAAATATAGIATGATATTAATATA 324	13 <i>30</i>
3244	GTTTTATTATATAATGCTTGTCTATTCAAGATTTGAGAACATTAATATGATACTGTCCACATATCCAA 331	12
	Ndel	35
3313	TATATTAAGTTTCATTTCTGTTCAAACATATGATAAGATGGTCAAATGATTATGAGTTTTGTTATTTAC 338	31
	3341	40
3382	Taqi Sau3ni Alui Reai CTGAAGAAAAGATAAGTGAGCTTCGAGTTTCTTGAAGGGTACGTGATCTTCATTTCTTGGCTAAAAGCGA 345	50
•	3402 3421 3405 3425	45
3451	ATATGACATCACCTAGAGAAAGCCGATAATAGTAAACTCTGTTCTTGGTTTTTGGTTTAATCAAACCGA 351	50
		55

		MspI HpaII A	DdeI luI		NdeI	MspI HpaII HinfI		3934
	3520 AC		1	AGCAAACATCGCAAACC	 ATATGTCAATTCG	TTAGATTCCCGGTT	TAA 3588	3334
5		3522 3	528 3529		3560	3576 3581 3581		
10			MapI HpaII					4003
	3589 G	TTGTAAA	CGGTATTTCATTT	GGTGAAAACCCTAGAAG	CCAGCCANCCTTT	TTTTAATCTAATTT	GCA 3657	
15			3598 3598	·				4072
20	3658 A	ACGAGAA	GTCACCACACCTC	CCACTAAAACCCTGAA	DdeI CCTTACTGAGAGA	HinfI HincII BetNI AGCAGAGNCANNAAJ	AGAA 3726	4141
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	3727 (AAATAAA :	ACCCGAAGATGAG	ACCACCACGTGCGGCGG	GACGTTCAGGGGA	ADADAADDADDDDD.	ATGR 3795	4210
30	2706 /	1	AluI	:GGCGGACGTTTTGGTGG	CGCCGTGGACGT	TTTGGTGGCGGCGG	AvaII TGGA 3864	
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40	3865	CCTTTGG	EcoRV IGGTGGATATCGT0	AvaII ACGAAGGACCTCCCAGT	GAAGTCATTGGT:	rcgtttactcttttc	DdeI CTTAG 3933	427
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2034	TaqI HinfI 	chaccacate CCCCC	_	HindIII AluI	- = ATTGATAAAGT	DdeI TCTCA 4002	
,,,,,	3937 3935			3976 3974		400C	5
4003	AluI GCTTTGAATGTG	XmnI RATGAACTGTTTCC	TGCTTATTAGTGTT	CCTTTGTTTTGAG	HinfI TTGAATCACTO	DdeI TCTTA 4071	10
	4004	4023			4059	4069	
4072	GCACTTTGTTA	HinfI GATTCATCTTTGTG 4085	TTTAAGTTAAAAGG	TAGAAACTTTGTG	SACTTGTCTCC	GTTATG 4140	
4141	Hincli I ACAAGGTTAACI	TTGTTGGTTATAA	:Agaagttgcgacci	TTCTCCATGCTT	STGAGGGTGAT	GCTGTG 4209	20
4210	A146 AvaII AluI	1 1	ni Cottacttoaatgoo	CCAATCTACTTG	Gaaaacaagac	ACAGAT 4278	25
12.10		4222 4231					30
4279	TGGGAAAGTTG	Hind Sau3AI <u>Al</u> ATGAGATCCAAGCT		II CI ECORI			35
			02 43 4314 4313 431	16 4321			
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The lambda CGN1-2 napin gene is that encoding the mRNA corresponding to the BE5 cDNA as determined by the exact match of their nucleotide sequences.

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An expression cassette was constructed from the 5'-end and 3'-end of the lambda CGN1-2 napin gene as follows in an analogous manner to the construction of pCGN944. The majority of the napin coding region of pCGN940 was deleted by digestion with Sall and religation to form pCGN1800. Single-stranded DNA from pCGN1800 was used in an in vitro mutagenesis reaction (Adelman et al., DNA (1983) 2:183-193) using the synthetic oligonucleotide 5'-

GCTTGTTCGCCATGGATACTTGTGTATGTTC-3'. This oligonucleotide inserted an EcoRV and an Ncol restriction site at the junction of the promoter region and the ATG start codon of the napin gene. An appropriate mutant was identified by hybridization to the oligonucleotide used for the mutagenesis and sequence analysis and named pCGN1801.

A 1.7 kb promoter fragment was subcloned from pCGN1801 by partial digestion with EcoRV and ligation to pCGN786 (a pCGN566 chloramphenicol based vector with synthetic linker described above in place of the normal polylinker) cut with EcoRI and blunted by filling in with DNA Polymerase I Klenow fragment to create pCGN1802. 3' sequences from the lambda CGN1-2 napin gene were added to Xhol/HindIII digested pCGN1802 to complete the cassette by ligation to pCGN941 digested with Xhol and HindIII. The resulting expression cassette, pCGN1803 contains 1.725 kb of napin promoter sequence, and 1.265 kb of napin 3' sequences with the unique cloning sites Sall, Bglll, Pstl and Xhol in between.

The ACP-precursor coding region from pCGN1Bcs is excised by double digestion with Xho1 and EcoRl and ligated to the cloning vector pUC18 previously digested with Sall and EcoRl. Transformation of ligated DNA into the appropriate E. coli host and screening using the penicillin-resistant, blue-white screening system of pUC vectors (Vieira and Messing, Gene (1982) 19:259-268) generates a plasmid containing a unique Dra3 site located downstream of the stop codon for the ACP-precursor coding region. Digestion with Dra3 followed by ligation with a Pstl linker generates a plasmid in which the coding region is cleanly excised as a Bglll to Pstl fragment, which is cloned into Pstl and Bglll sites of pCGN1803 with the proper orientation of the ACP-precursor coding region to the napin promoter and terminator parts of the embryo-specific expression information encoded by pCGN1803. The resulting chimeric gene is then introduced into a binary vector and transferred to Agrobacterium for cocultivation with Brassica napus hypocotyl segments employing the same conditions employed for the spinach ACP chimeric gene in pCGN946.

In accordance with the subject invention, sequences coding for functional acyl carrier protein, particularly plant acyl carrier protein, are provided, which can be used as probes for detecting the presence of acyl carrier protein genes, for screening libraries from plants and bacteria, either genomic or cDNA, for use in assays for detecting the presence of acyl carrier protein genes, and the like. In addition, the coding sequence can be used in the preparation of expression constructs, where the coding sequence is combined with transcriptional and translational initiation and termination regulatory regions for expression in an appropriate host in which the regulatory regions are functional. A particular interest is the use of the ACP coding sequence in conjunction with a transcription initiation region which is functional in a plant and particularly is regulated so as to provide for expression in seed. In this manner, the production of seed oil can be enhanced and, as appropriate, the fatty acid composition modulated.

All publications and patent applications mentioned in this specification are indicative of the level of skill of those skilled in the art to which this invention pertains. All publications and patent applications are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

45 Claims

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- 1. A cDNA sequence coding for plant acyl carrier protein.
- 2. A cDNA sequence according to Claim 1, wherein said plant is spinach.
- 3. A cDNA sequence according to Claim 1, wherein said plant is Brassica.
- 4. A cDNA sequence according to Claim 3, wherein said Brassica is campestris.
- 5. A cDNA sequence according to Claim 3, wherein said Brassica is napus
- 6. A DNA construct comprising an open reading frame coding for a plant acyl carrier protein joined to and under transcriptional and translational initiation and termination regulatory regions functional in a cellular host, wherein at least one of said regulatory regions is other than the wild-type region of said open reading frame.
- 7. A DNA construct according to Claim 6, wherein said open reading frame encodes for spinach or Brassica acyl carrier protein.
 - 8. A DNA construct according to Claim 6, wherein said regulatory regions are functional in a plant host.
- 9. A DNA construct according to Claim 8, wherein said transcriptional initiation region is regulated during cell differentiation.
- 10. A DNA construct according to Claim 9, wherein said regulation provides for transcription in the developing embryo.
- 11. A DNA construct according to Claim 10, wherein said transcriptional initiation regulatory region is a napin regulatory region.
- 12. A Ti- or Ri-plasmid comprising a DNA construct according to any of Claims 6 to 11.

13. A plant cell comprising a DNA construct according to any of Claims 6 to 11. 14. A plant cell according to Claim 13, wherein said plant cell is an embryonic cell and part of a seed. 15. A plant cell according to Claim 14, wherein said plant is a <u>Brassica</u> plant. 16. A method for enhancing the formation of vegetable seed oils which comprises, growing a plant from plantlet to seed, wherein the cells of said plant comprise a construct according to Claim 8, and harvesting the resulting seed containing an enhanced amount of vegetable oil. 17. A method according to Claim 16, wherein said plant is a Brassica plant.	5
18. A method according to any of Claims 16 or 17, wherein said transcriptional initiation regulatory region is a napin region.	10
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